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Patents ADP number (if you know it)

0744519001

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Tagging and Recovery of Elements Associated with Target Molecules

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91001

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Tagging and Recovery of Elements Associated
with Target Molecules

The present invention relates to a new method for
5 identifying elements associated with target molecules.

Many genes and gene clusters are controlled by known (or
unknown) distant regulatory elements that are necessary
for high-level expression. Identification of these
10 regulatory elements is an expensive and time-consuming
process. Previous attempts to identify such distant
regulatory elements have used a number of different
methods, but most directly by scanning large genomic
regions for DNase I hypersensitivity sites, followed by
15 functional analysis of those regions linked to reporter
genes in transgenic mice. This method of identification
will clearly take a very long time.

The beta-globin locus is the prototypical gene cluster
20 regulated by distant regulatory elements; the search for
the beta-globin regulatory elements took approximately 10
years. Experiments designed to locate the beta-globin
gene regulatory elements began in the late 1970s. In the
early 1980s data arose that suggested distant elements
25 were involved. A thalassemia patient was studied whose
genome contained an intact beta-globin gene but a large
deletion upstream of the gene. This lead to the
conclusion that a distant upstream element must be
involved in the regulation of the gene (Kioussis et al.,
30 1983). Indeed, transgenes containing the beta-globin gene
alone achieve only very low levels of expression at best
(Townes et al., 1985). In 1985 a series of DNase I

hypersensitive sites were mapped 40-60 Kb upstream of the beta-globin gene (Tuan et al., 1985). In 1987 it was finally shown that this hypersensitive site region, collectively known as the locus control region (LCR), was 5 sufficient to induce high-level, position-independent, copy number- dependent gene expression when linked to the beta-globin gene (Grosveld et al., 1987). Defects in human beta-globin gene expression, or hemoglobinopathies, are the most common genetic diseases worldwide. The ability to 10 induce high-level expression of an artificially introduced beta-globin gene is therefore of significant therapeutic use. In addition, the ability to locate control regions of other genes is clearly desirable.

15 Chromatin conformation capture (3C; Decker et al 2002) has previously been used to determine the conformation of a yeast chromosome. However, many technical problems arise when trying to apply this method to higher eukaryotes, not least because the mammalian genome is approximately 200 20 times the size of a yeast genome. The 3C has several disadvantages: 3C does not enable recovery of *in situ* labelled molecules, nor does 3C give a very high degree of resolution. In addition, other disadvantages of the 3C technique result because this technique allows only an 25 average conformation of a chromosome to be calculated; this means that if all the cells used in the technique are not homogeneous or the molecular conformation is dynamic, specific interactions may be overlooked. Further, the 3C technique does not provide a method for determining which 30 proteins or other molecules are associated with the genome.

Fluorescence *in situ* hybridisation (FISH) is a previously known technique which uses hapten-labelled nucleotide

probes followed by anti-hapten antibodies conjugated to fluorophores, to determine the site of an actively transcribed gene using the antibody's ability to specifically bind to the hapten. Covalent tag deposition 5 has commonly been used to enhance the signals obtained using the above technique. Kits enabling performance of covalent tag deposition to enhance signals are obtainable from NEN Dupont and are called TSA™ (Tyramide Signal Amplification™). However, this technique has not provided 10 means for purifying molecular complexes from specific sites or in the immediate vicinity of specific sites in or on cells. Neither FISH nor TSA allow for detection (and thus identification) of, for example, the interaction of distant regulatory elements with an actively transcribed 15 gene. There is no technique presently available to use for detecting (and thus identifying) the interaction of distant regulatory elements with an actively transcribed gene during the time of transcription.

20 Techniques are known which can be used for identification and analysis of proteins involved in protein complexes. ImmunoPrecipitation (IP) is most commonly used to "pull down" proteins associated in a complex with a target protein(s). However no techniques exist to analyse, for 25 instance, molecules or complexes which are only involved in "loose" functional interactions with another complex or which only function in the vicinity of another protein.

According to the present invention there is provided, a 30 method for identifying elements associated with target molecules comprising the steps of:

(a) providing a probe capable of binding specifically to a target molecule, the probe associated with an enzyme;

(b) adding a tag capable of being activated by the enzyme such that it can attach to elements in the vicinity of the enzyme; and

(c) isolating elements having the tag attached thereto.

5

The target molecules may include RNA molecules, DNA molecules, proteins or peptides, lipids, or other, artificial compounds.

10 When the target is RNA, the elements which may be associated with these target molecules and which may be identified (or whose mode of action can be understood) by using the technique of the present invention include: distant regulatory elements (i.e. DNA elements via their chromatin protein association) that are in proximity to the RNA of an actively transcribed gene; RNA binding proteins such as those involved in RNA processing or stabilization/regulation/etc; proteins and protein complexes which facilitate the interactions between regulatory elements and a gene; proteins and protein complexes involved in the activation of genes; proteins and protein complexes involved in the regulation of chromatin structure in and around active genes; and transcription factors.

15

25

When the target is DNA, the elements which may be associated with these target molecules and which may be identified (or whose mode of action can be understood) by using the technique of the present invention include: distant regulatory elements (i.e. DNA elements via their chromatin protein association) that are in proximity to the targeted DNA; other DNA elements in proximity to the targeted DNA, which may be for example, engaged in

functional interactions with the target sequence (e.g. boundaries, insulators, structural or architectural interactions); analysis of higher order chromatin structure, for example the analysis of tertiary chromatin interactions (chromatin folding); mapping chromatin interactions in entire loci or whole genomes (with the aid of high throughput technology); protein/protein complexes involved in regulation of gene expression or the control of chromatin structure.

When the target is protein, the elements which may be associated with these target molecules and which may be identified (or whose mode of action can be understood) by using the technique of the present invention include: DNA elements in proximity to a protein; RNA molecules in proximity to a protein; or other proteins/protein complexes bound to, or in the vicinity of a targeted protein (e.g. identifying other protein components of the LCR-beta-globin gene complex at different stages of development, or identifying the *in-vivo* ligands of a specific receptor- or vice versa).

When the target is lipid, the elements which may be associated with these target molecules and which may be identified (or whose mode of action can be understood) by using the technique of the present invention include: DNA elements in proximity to a lipid or artificial compound RNA molecules in proximity to a lipid or artificial compound; or proteins/protein complexes bound to, or in the vicinity of a targeted lipid or artificial compound. The probe usable in the present invention may be a DNA probe, an RNA probe or an antibody specific for a protein, lipid or other molecule.

The probes used can be associated with the enzyme through antibody/enzyme conjugates, or enzyme/target molecule fusion.

5

The method by which the enzyme may be targeted to a specific molecule may be varied depending on the molecule to be targeted. For example, using a labelled probe specific for a DNA molecule, using immuno-histochemistry, 10 or using a fusion of a protein (or other molecule of interest) and the enzyme. Preferably antibody/enzyme conjugates may be used. In one preferred embodiment, when the target molecule is RNA, a hapten-labelled probe specific to the intron of an active gene can be added, 15 followed by addition of a hapten-specific Fab fragment/enzyme conjugate. One hapten which may be used is digoxigenin (DIG); others include biotin, dinitrophenol and FITC.

20 An enzyme which may be used in the present invention is Horse Radish Peroxidase. This enzyme can be used in combination with a tyramide molecule such as biotin-tyramide, dinitrophenol-tyramide or FITC-tyramide. Another enzyme/TAG combination is ubiquitin-conjugating enzyme, 25 with ubiquitin as a tag.

30 Protein kinase could also be used as the enzyme (there are several with varied specificities) with phosphate as a tag. In this example a kinase which is able to add a phosphate to a nucleosomal protein (if looking for chromatin tagging) or other protein of interest should be used. Antibodies against the specifically modified epitope of the particular amino acid residue receiving the

phosphate could be used to target and isolate the tagged elements.

DNA Adenine Methyltransferase (DAM) is another enzyme which could be used, with a methyl group as the tag. In a slight variation of the procedure, instead of using a tag to pull out the labelled material one could use a restriction enzyme that will cut only DNA which is specifically methylated by DAM. DAM adds a methyl group to the adenine in the sequence GATC. This methylated site can only be cut by the DNA restriction endonuclease DpnI. DAM is normally only found in bacteria such as E.coli so it could be used in eukaryotic cells without any interference from endogenous methyltransferases which only methylate other sequence combinations. With this method no affinity chromatography is required. DNA can simply be purified from the DAM treated cells, cut with DpnI and small DNA fragments released from the mixture of genomic DNA can be isolated. The small sites released by DpnI digestion can then be labelled with radioisotopes, etc., and used for diagnostic hybridization to, for example, a microarray (van Steensel et al 2001)

Other enzyme/tag combinations could be used: any enzyme which can activate a tag molecule to deposit onto another molecule, for example protein, DNA, RNA, lipid etc in a manner such that the tagged product can then be isolated by whatever means (eg. affinity chromatography or immunoprecipitation) can be used in this technique.

30

Before separation, the molecules which have been tagged can be disrupted into smaller fragments using, for example, sonication, enzymatic cleaving, shearing with a

French Press or small bore syringe, or any other method which achieves such a result.

Analysis of the DNA obtained using the above method can be
5 used to identify any regulatory elements which were in proximity to the active gene, because these elements become labelled with the tag, due to their proximity to the site HRP activity. The DNA can then be analysed by a number of quantitative techniques, for example
10 Quantitative PCR (for example Real-Time PCR (Wittwer et al., 1997)) or semi-quantitative PCR, slot blot or microarray (Granjeaud et al., 1999), among others. Real-time PCR analysis allows scanning, high-throughput, high resolution analysis of any gene locus for hundreds or
15 thousands of kilobases in either direction.

An embodiment of the present invention will now be described in more detail, by way of example, with reference to the drawings, in which:

20

Figure 1 is a schematic diagram showing a method of the present invention

25

Figure 2 is a schematic diagram showing the mouse beta-globin locus and locus control region (LCR) and two models of LCR action; and

30

Figure 3 is a schematic diagram showing the hypothesised interaction of the mouse beta-globin locus and locus control region (LCR), as a result of performing the present invention.

Figure 4 is a graph showing the results of quantitative real-time PCR analyses of β maj-directed RNA TRAP (top curve) showing various sequences in the β globin locus and neighbouring olfactory receptor gene locus using the primer pairs shown.

Figure 5 is a slot blot analysis of β maj directed RNA TRAP.

10 Figure 6 is a diagram showing the results of a double label RNA FISH using β maj and β min intron probes.

Figure 7 is a graph showing the results of β min-directed RNA TRAP assaying various sequences in the β globin locus.

Many genes and gene clusters are thought to be regulated by distant regulatory elements, which may be located tens to hundreds of kilobases away. The best characterised example of a distant element regulating a cluster of genes is the beta-globin locus control region (LCR), shown in Figure 1. The LCR 7 consists of a series of Dnase I hypersensitive sites (HS) (1 to 6). At the core of each HS is a 200-300 base-pair (bp) region packed with multiple binding sites for ubiquitous and tissue-specific factors. The LCR 7 is absolutely required for high- level transcriptional-activation of all the beta-globin genes. Two models have been proposed to explain the action of the LCR, although no direct proof exists for either mode of action. These models are shown in Figure 2. The first model 8 proposes that the LCR works at a distance and hypothesises that the LCR creates a large region of open

chromatin surrounding the genes and recruits and sends factors necessary for gene activity along the chromatin. The second model 9 proposes that the LCR physically contacts the gene(s) through long range chromatin interactions, essentially looping out the intervening sequences and activating transcription directly.

To determine if an actively transcribed beta-globin gene is in direct physical contact with the distant (40Kb) LCR 10 7 *in vivo*, the following technique can be used (see Figure 2). Firstly, fetal liver 10, the main site of erythropoiesis in the developing foetus, is taken and disrupted, and the cells spread in a monolayer on a slide 11, prior to cross-linking with formaldehyde. *In-situ* 15 hybridization is performed using a digoxigenin (DIG)-labelled oligonucleotide probe 12, specific for the intron of the mouse beta-major globin gene. The enzyme Horse Radish Peroxidase (HRP) is then targeted to an RNA molecule using an anti-DIG antibody conjugated to Horse 20 Radish Peroxidase (HRP) 13, thus pinpointing HRP enzyme activity to the site of the actively transcribed gene.

Next, biotin-tyramide 14 is added as a molecular tag; it is activated by the HRP to cause it to covalently attach 25 to electron-dense amino-acids in the immediate vicinity. After the tag is covalently attached 15, the cells are sonicated to give small, soluble chromatin fragments 16 having an average DNA size of 400bp. The biotinylated chromatin is then purified using streptavidin-agarose 30 affinity-chromatography 17, cross-links are reversed and the DNA is purified. Multiple amplicons across the locus can then be analysed 18 using quantitative or semi-quantitative PCR and/or slot blotting.

By using the above technique on the mouse beta-globin gene locus, it was found that the LCR and active beta-major gene are in significant proximity *in vivo*; HS2 2 appears 5 to be in intimate contact with the beta-major gene.

EXAMPLES

Example 1 RNA FISH-TRAP

10 E14.5d fetal livers from Balb/c mice, in which only the adult-type b-maj and b-min genes are expressed, were disrupted in ice-cold PBS. The cells were spread on poly-L-lysine coated slides and fixed in 4% formaldehyde, 5% acetic acid for 18 minutes at room temperature. Subsequent 15 slide-washing, permeabilization, probe-hybridisation, and post-hybridisation washing were performed as described in Gribnau, J. et al. (1998); the probes used being directed to intron 2 near the 3' end of the mouse b-maj globin primary transcript. Endogenous peroxidases were quenched 20 in 0.5% H₂O₂ (in PBS) for 10 minutes followed by washing (5min) in TST (Tris, saline, Tween; 100mM Tris pH7.5, 150mM NaCl, 0.05% Tween 20) and blocking as described. Slides were then incubated with 1:100 dilution of anti-DIG 25 Fab fragment/HRP conjugate for 45 minutes at room temperature in a humidified chamber, washed twice (5 min each) in TST and then incubated for 1 minute with 1:150 biotin-tyramide (NEN) under coverslips at room temp. The slides were then quenched again in 0.5% H₂O₂ (in PBS) for 10 minutes, washed twice in TST (5min) and transferred to 30 PBS ready for scraping. One of the slides was stained with an Avidin/Texas Red conjugate for 45 minutes at room temperature. This slide was then washed, dehydrated,

mounted and visualised as described in Gribnau, J. et al. (1998).

Cells were scraped from the remaining slides; typically 5 approximately 25 million cells were recovered. The cells were spun down at 2900g for 25 minutes, resuspended in 2M NaCl, 5M Urea, 10mM EDTA, and sonicated for 200 seconds on ice (eight 25-second bursts with 1.5 minutes between bursts) using a Microson Ultrasonic Cell Disruptor set at 10 level 5. Crude chromatin was centrifuged for 15 minutes at 10,000g, the supernatant containing the soluble chromatin was removed and the insoluble pellet was resuspended in 2M NaCl, 5M Urea, 10mM EDTA, and sonicated again. The suspension was centrifuged again and the two soluble 15 fractions were combined and dialysed overnight at 4°C against PBS. This method routinely yielded chromatin fragments with an average DNA size of around 400bp.

10% of the soluble chromatin was set aside as the input 20 and the rest was passed over a streptavidin-agarose (Molecular Probes) affinity column. After binding, the column was washed with 3X700µl PBS, 2X500µl TSE 150 (20mM Tris pH8.0, 1% Triton, 0.1% SDS, 2mM EDTA, 150mM NaCl), 2X500µl TSE 500 (20mM Tris pH8.0, 1% Triton, 0.1% SDS, 2mM EDTA, 500mM NaCl), and 3X700µl PBS. The beads were then 25 removed from the column, formaldehyde cross-links reversed and protein components digested by overnight incubation at 65°C with 200µg/ml proteinase K while shaking vigorously. The samples were treated with 20µg/ml RNase A for 30min at 30 37°C, 200µg/ml proteinase K for 5 hours at 37°C, phenol-extracted and ethanol-precipitated using 20mg/ml glycogen as carrier. DNA from the input (IP) fraction was quantified using a standard spectrophotometer. DNA concentration of the affinity purified (AP) fraction was

measured by picogreen quantification using IP as a standard.

Example 2 REAL-TIME PCR

5 Real-time PCR was performed with an ABI PRISM 7700 sequence detector using 2X SYBR green PCR master mix (Applied biosystems). For each primer pair a standard curve was generated using 30ng, 5ng, and 1ng of IP which was then used to quantify the enrichment of 1ng of AP (all 10 reactions were performed in duplicate). All PCR products were run on a 2% agarose gel to ensure all reactions gave a single product.

15 Enrichment of various sequences across the β -globin locus and also across the neighbouring olfactory receptor gene (*org*), were measured using quantitative real-time PCR. The measurements showed a 20-fold peak of enrichment near the transcription termination site of the β -maj gene, consistent with the position of the probes (Figure 4; top 20 curve). Enrichment dropped off sharply upstream of the β -maj gene for over 25kb in the area of the developmentally silenced γ and β H1 genes, which are only slightly increased over background.

25 Strikingly, a peak of enrichment was observed over HS2, and to a lesser extent HS1 and HS3 of the LCR. This indicates these sites are in close association with the active gene.

30 The fact that other HS in the LCR (HS4, 5 and 6) and the downstream 3'HS1 (which is closer in base pairs to the β maj gene than HS2) are not significantly enriched

suggests they are outside the area of labelling and therefore not intimately associated with the active β _{maj} gene. Moreover, the low level of enrichment of these sites shows that there is no preferential labelling of areas of hypersensitive or open chromatin. To completely discount the possibility that these results were caused by a bias of biotin deposition in certain areas (e.g. open or hyperacetylated chromatin) a control random TRAP experiment was designed and performed. By omitting the intron probe during the FISH-stage, biotin deposition becomes random across the genome and therefore any bias for certain sequences would become apparent in the analysis of the AP material. Fig. 4, lower curve, shows no preferential selection for any of the sequences in the globin locus, thus verifying that enrichment of HS2 in the β _{maj}-directed TRAP experiment is due to proximity to the active β _{maj} gene and is not a chromatin bias. Repetition of the β _{maj} RNA TRAP assay three times obtained similar results. DNA from one of the β _{maj} RNA TRAP assays was analyzed by slot blot with multiple probes yielding similar results (Figure 5). The data of this experiment provide the first direct evidence that a distal enhancer is held in significant physical proximity to an active gene that it regulates *in vivo*.

To distinguish between a co-transcriptional model in which both genes share the LCR simultaneously or an alternating model in which the LCR is involved exclusively with a single active gene. RNA-TRAP was repeated using intron probes to the β _{min} gene located approximately 15 kb downstream of β _{maj}. The results of this showed that HS2 is highly enriched in the β _{min}-directed AP chromatin, indicating it is tightly associated with the active β _{min} gene (Figure 7). In addition, HS4 of the LCR was

significantly enriched over background levels and when compared to HS1, 3, 5 and 6 of the LCR. The high level of enrichment of HS2 in both the β min and β maj directed RNA-TRAP assays indicates it is tightly associated with the 5 active gene for most of the time primary transcript is present. The fact that β maj-TRAP does not bring down the β min gene and vice versa indicates the two genes are not closely associated.

10 There are many applications for the technique of the present invention, which can be performed in vivo, ex vivo, or in vitro. One example of such a use is in transgenic animal technology: transgenic animals are presently being used by a number of laboratory around the 15 world as bioreactors to produce large amounts of proteins of interest. The most commonly used method is to express the protein of interest in milk under control of a highly expressed milk protein gene promoter. Most transgenic animals created with such a construct would not express 20 the protein or express it at very low levels making them unusable. Some transgenic animals may, by virtue of position effects at the site of integration of the construct, express larger amounts of the protein of interest. The addition of milk protein gene LCR-like 25 sequences to the expression construct would increase the number of transgenic animals which express the gene to 100% and increase the average level of expression in every animal. This would significantly decrease the cost of production and greatly increase the yield.

30

When RNA is the target molecule, the method of the present invention labels only the cells in the population that are actively transcribing the gene of interest. The advantage of this is specifically interacting sequences are highly

enriched upon affinity chromatography, whether the population is heterogeneous or the interaction is dynamic (Wijgerde et al., 1995). Another advantage of the present invention when RNA is the target molecule is this

5 technique can detect (and thus identify) the interaction of distant regulatory elements with an actively transcribed gene during the time of transcription. There

is no other technique we know of which can be used for this purpose. This technique can specifically label and

10 recover proteins at the site of transcription in a dynamic or heterogeneous population of cells and identify specific interactions.

Another advantage of the present invention which results

15 whatever the target molecule is, is the possibility of labelling and recovering complexes in the vicinity of a target complex (as opposed to molecules which are in direct interaction). The resultant enriched proteins could be analysed by a number of protein chemistry

20 techniques such as Western blotting, Mass Spectroscopy, fractionation, purification, polyacrylamide gel electrophoresis, etc.

The present invention provides a relatively easy and rapid

25 method which can detect interactions between an actively transcribed gene and distant regulatory element(s). The technique can also be used to identify any sequence element involved in an interaction with any other target sequence *in vivo* by virtue of their proximity.

30

The present invention provides a new way to identify the regulatory elements involved in the activation of genes in a rapid and relatively inexpensive way. It has also been

used to address the question of how LCRs or enhancer elements function and in fact has provided the first direct evidence that the LCR functions by physically interacting with an actively transcribed gene in the beta-
5 globin locus.

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CLAIMS

1. A method for identifying elements associated with target molecules comprising the steps of:

5

(a) providing a probe capable of binding specifically to a target molecule, the probe associated with an enzyme;

10

(b) adding a tag capable of being activated by the enzyme such that it can attach to elements in the vicinity of the enzyme; and

15

(c) isolating elements having the tag attached thereto.

20

1. A method as claimed in claim 1 in which the target molecule is selected from the group consisting of RNA molecules, DNA molecules, proteins or peptides, lipids, or other, artificial compounds.

25

2. A method as claimed in claim 1 or 2 in which the elements which may be associated with the target molecules include distant regulatory elements, RNA, DNA, proteins and protein complexes, transcription factors, or in-vivo ligands of a specific receptor.

30

3. A method as claimed in any preceding claim in which the probe is selected from the group consisting of DNA probe, an RNA probe or an antibody specific for a protein, lipid or other molecule.

4. A method according to claim 4 in which the probe is associated with the enzyme through an antibody/enzyme conjugate, or enzyme/target molecule fusion.
- 5.
- 6.
- 10 5. The method according to any preceding claim in which the enzyme is targeted to RNA using a hapten-labelled probe specific to the RNA of an intron of an active gene, and then a hapten-specific Fab fragment/enzyme conjugate is added.
- 15 6. The method according to any preceding claim in which the hapten is digoxigenin, biotin, dinitrophenol or FITC.
- 7.
- 20 8. The method according to any preceding claim in which elements are isolated using affinity chromatography or ImmunoPrecipitation.
- 9.
- 25 25. A method for identifying elements of chromatin associated with transcribing RNA comprising the steps of:
 - (a) providing a hapten-labelled probe capable of binding specifically to RNA of a gene,

(b) providing an antibody conjugated with the enzyme horse-radish peroxidase, the antibody specific for the hapten;

5 (c) adding biotin-tyramide such that it can attach to elements in the vicinity of the enzyme;

(d) disrupting the chromatin

10 (e) isolating elements of chromatin having biotin attached thereto using affinity chromatography and purifying the elements.

10. The method of claim 9 in which the chromatin is disrupted using sonication, enzymatic cleaving, or shearing with a French Press or small bore syringe.

11. The method according to any of claims 9 to 10s in which the hapten is digoxigenin.

20 12. Elements isolated by the method of any preceding claim.

25 13. Analysis of DNA obtained using the method according to any preceding using Quantitative Real-Time PCR, slot blot or microarray.

14. A method for identifying DNA associated with target molecules comprising the steps of:

(a) providing a probe capable of binding specifically to a target molecule, the probe associated with an DNA Adenine Methyltransferase;

5 (b) adding a restriction enzyme that will cut only DNA specifically methylated by DAM;

(c) isolating DNA cut by the restriction enzyme

10 (d) identifying the isolated DNA.

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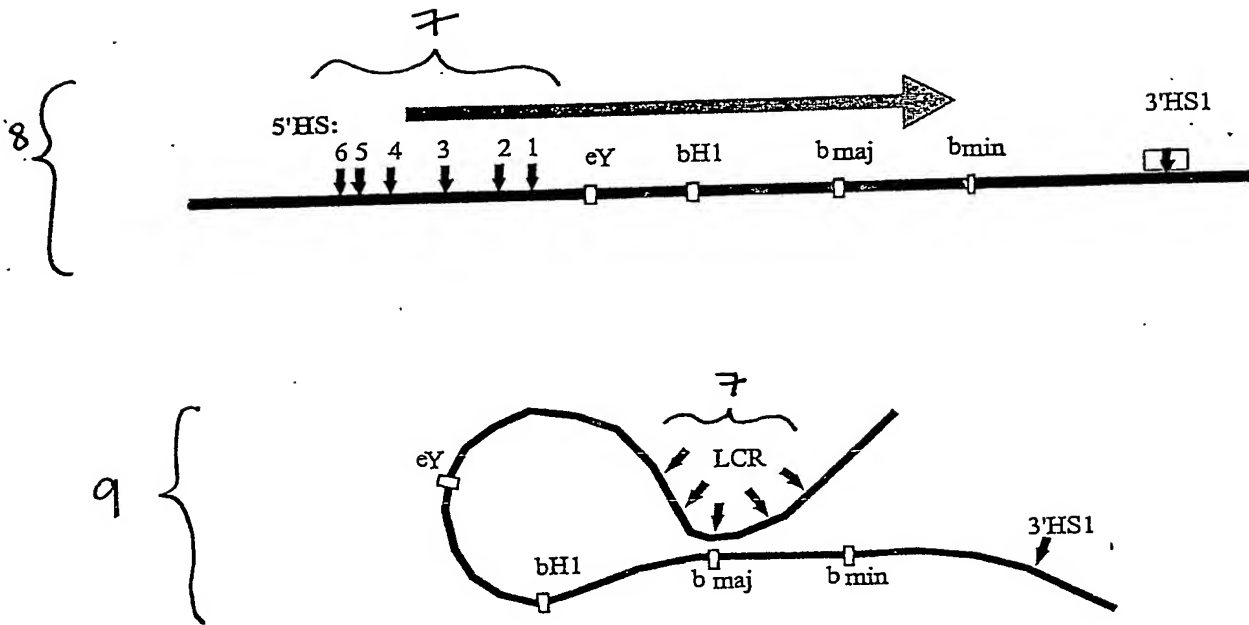


FIGURE 1

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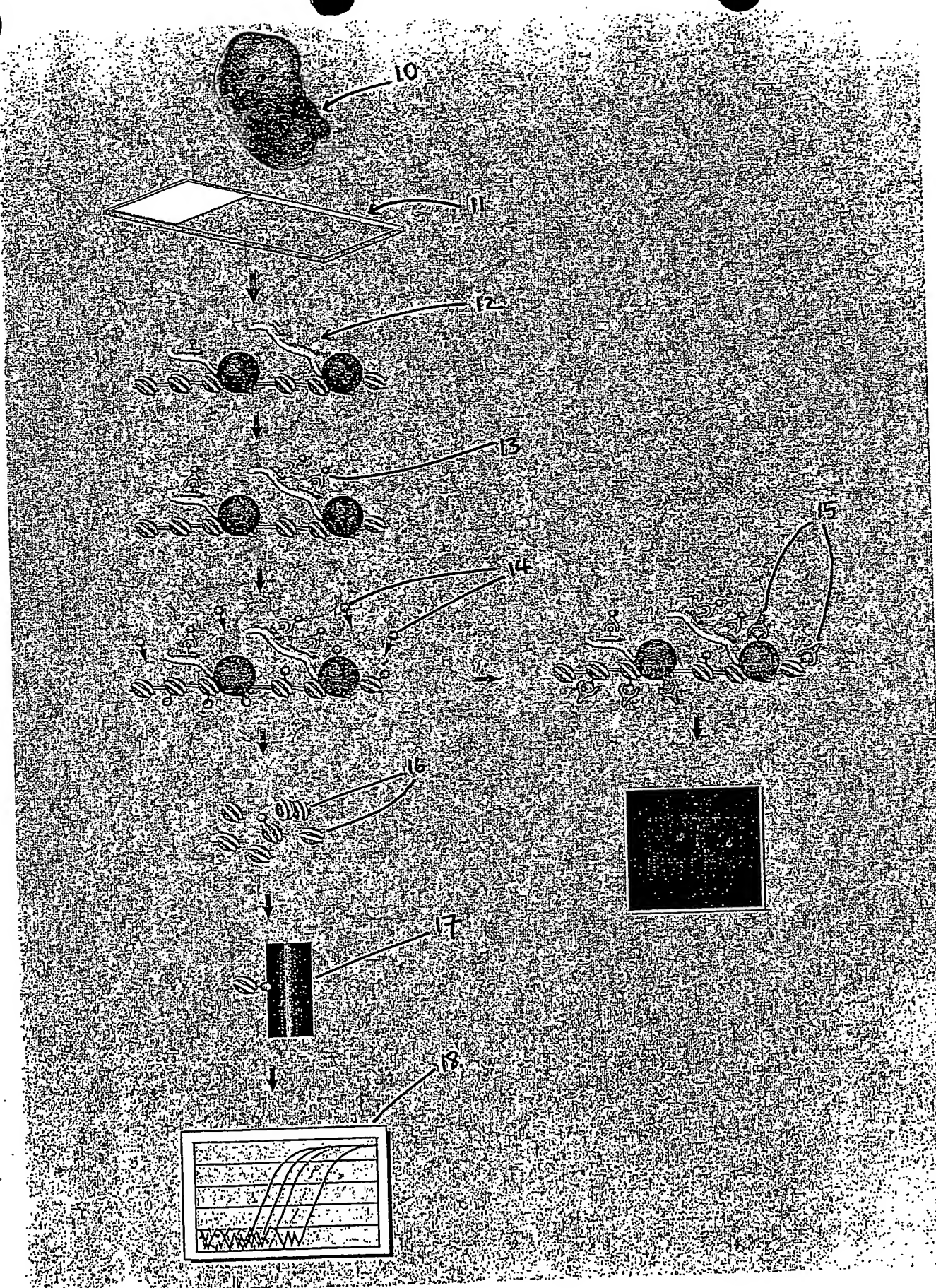


FIGURE 2

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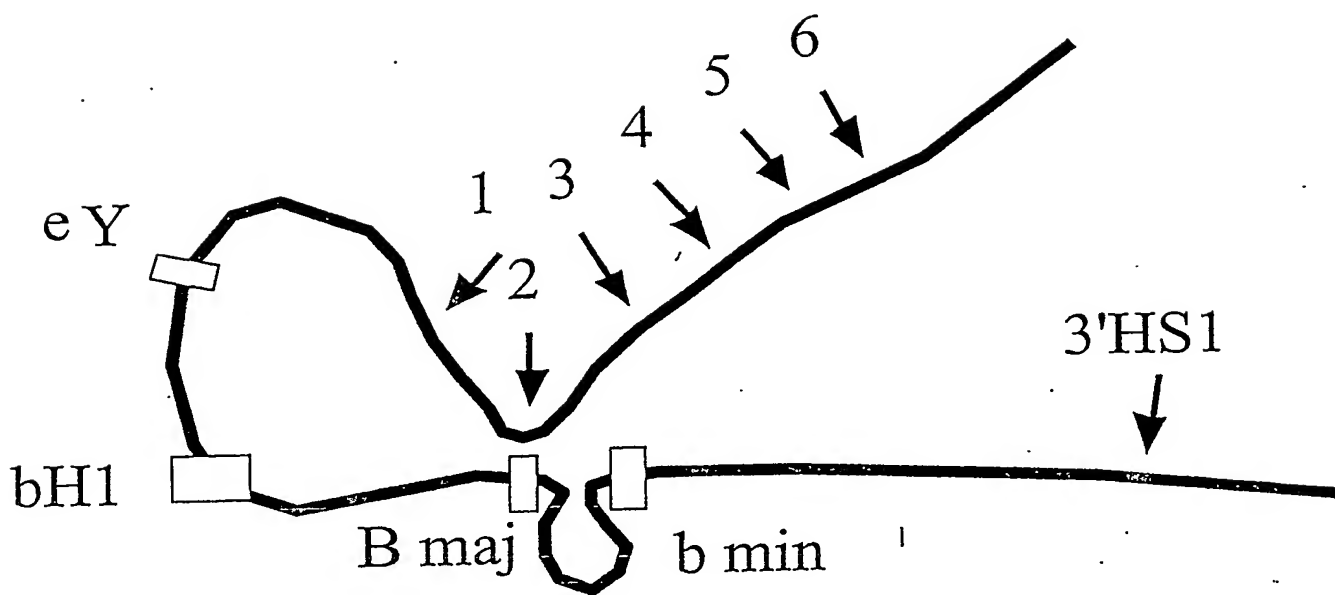


FIGURE 3

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FIGURE 4

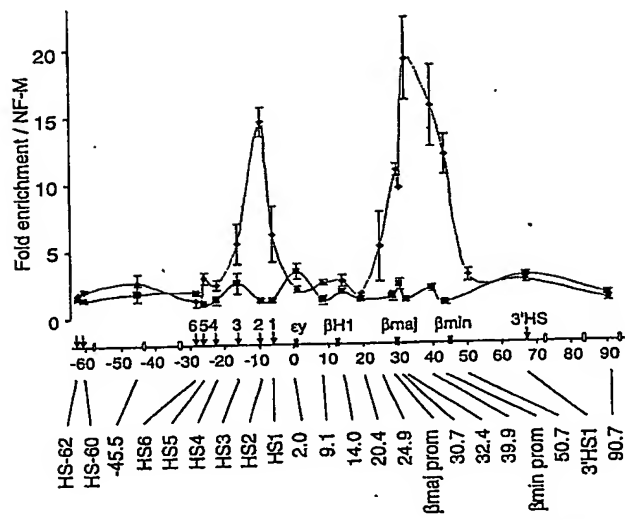


FIGURE 5

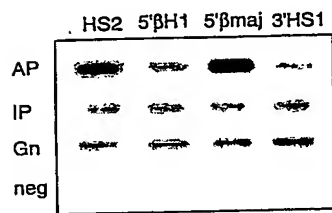
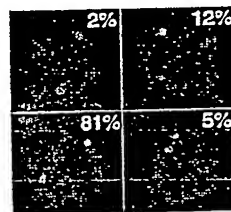


FIGURE 6



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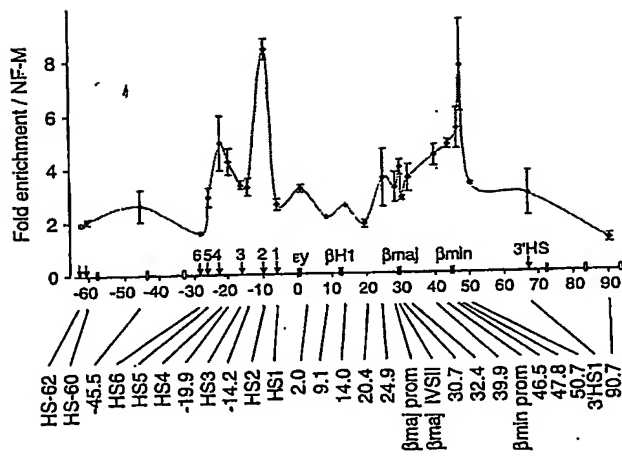
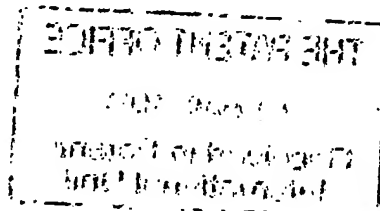


FIGURE 7



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